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(54) Title: A NOVEL β1->6 N-ACETYLGLUCOSAMINYLTRANSFERASE, ITS ACCEPTOR MOLECULE, LEUKO-SIALIN, AND A METHOD FOR CLONING PROTEINS HAVING ENZYMATIC ACTIVITY

(57) Abstract

The present invention provides a novel β 1->6 N-acetylglucosaminyltransferase, which forms core 2 oligosaccharide structures in O-glycans, and a novel acceptor molecule, leukosialin, CD43, for core 2 β 1->6 N-acetylglucosaminyltransferase activity. The amino acid sequences and nucleic acid sequences encoding these molecules, as well as active fragments thereof, also are disclosed. A method for isolating nucleic acid sequences encoding proteins having enzymatic activity is disclosed, using CHO cells that support replication of plasmid vectors having a polyoma virus origin of replication. A method to obtain a suitable cell line that expresses an acceptor molecule also is disclosed.

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A NOVEL B1+6 N-ACETYLGLUCOSAMINYLTRANSFERASE,

ITS ACCEPTOR MOLECULE, LEUKOSIALIN, AND
A METHOD FOR CLONING PROTEINS HAVING ENZYMATIC ACTIVITY

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BACKGROUND OF THE INVENTION

FIELD OF THE INVENTION

This invention relates generally to the fields of 10 biochemistry and molecular biology and more specifically to a novel human enzyme, UDP-GlcNAc:Galß1→3GalNAc (GlcNAc to GalNAc) β1→6 N-acetylglucosaminyltransferase (core 2 β1→6 N-acetylglucosaminyltransferase; C2GnT), and to a novel 15 acceptor molecule, leukosialin, CD43, for core 2 $B1\rightarrow6$ Ninvention acetylglucosaminyltransferase action. The additionally relates to DNA sequences encoding core 2 81+6 N-acetylglucosaminyltransferase and leukosialin, to vectors containing a C2GnT DNA sequence or a leukosialin DNA 20 sequence, to recombinant host cells transformed with such vectors and to a method of transient expression cloning in CHO cells for identifying and isolating DNA sequences encoding specific proteins, using CHO cells expressing a suitable acceptor molecule.

BACKGROUND INFORMATION

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Most O-glycosidic oligosaccharides in mammalian glycoproteins are linked via N-acetylgalactosamine to the hydroxyl groups of serine or threonine. These O-glycans can be classified into 4 different groups depending on the nature of the core portion of the oligosaccharides (see Fig. 1). Although less well studied than N-glycans, O-glycans likely have important biological functions. Indeed, the presence of O-linked oligosaccharides with the

core 2 branch, Galß1→3(GlcNAcß1→6)GalNAc, has been demonstrated in many biological processes.

Piller et al., <u>J. Biol. Chem</u> 263:15146-15150 (1988) reported that human T-cell activation is associated 5 with the conversion of core 1-based tetrasaccharides to core 2-based hexasaccharides on leukosialin, a major sialoglycoprotein present on human T lymphocytes (see also A similar increase in hexasaccharides was observed in peripheral blood lymphocytes of patients suffering from T-cell leukemias (Saitoh et al., Blood 77:1491-1499 (1991)), myelogenous leukemias (Brockhausen et al., Cancer Res. 51:1257-1263 (1991)) and immunodeficiemcy due to AIDS and the Wiskott-Aldrich syndrome (Piller et al., J. Exp. Med. 173:1501-1510 (1991)). patients' 15 lymphocytes, changes in the hexasaccharides were caused by increased activity of either UDP-GlcNAc:Galß1→3GalNAc (GlcNAc to GalNAc) acetylglucosaminyltransferase (EC2.4.1.102) or core 2 \mathfrak{R}_{\bullet} N-acetylglucosaminyltransferase (Williams et al., <u>J. Biol.</u> 20 <u>Chem.</u> 255:11253-11261 (1980)). Increased activity of core 2 $\mathfrak{S}1 \rightarrow 6$ N-acetylglucosaminyltransferase also was observed in metastatic murine tumor cell lines as compared to their parental, non-metastatic counterparts (Yousefi et al., J. Biol. Chem. 266:1772-1782 (1991)).

Increased complexity of the attached oligosaccharides increases the molecular weight of the glycoprotein. For example, leukosialin containing hexasaccharides has a molecular weight of ~135kDa, whereas leukosialin containing tetrasaccharides has a molecular weight of ~105kDa (Carlsson et al., J. Biol. Chem. 261:12779-12786 and 12787-12795 (1986)).

Fox et al., <u>J. Immunol.</u> 131:762-767 (1983) raised a monoclonal antibody, T305, against human T-lymphocytic leukemia cells. Sportsman et al., <u>J. Immunol.</u> 135:158-164

(1985) reported T305 binding was abolished by neuraminidase treatment, suggesting T305 binds to hexasaccharides. T305 specifically reacts with the high molecular weight form of leukosialin (Saitoh et al., supra, (1991)).

5 Previous studies indicated acetyllactosamine repeats extend almost exclusively from the branch formed by the core 2 B1→6 Nacetylglucosaminyltransferase (Fukuda et al., J. Biol. <u>Chem.</u> 261:12796-12806 (1986)). Consistent with these 10 results, Yousefi et al., supra, (1991) demonstrated that the core 2 enzyme in metastatic tumor cells regulates the level of poly-N-acetyllactosamine synthesis in O-linked oligosaccharides.

Poly-N-acetyllactosamines are subject to a variety of modifications, including the formation of the sialyl Le^x, NeuNAcα2→3Galß1→4(Fucα1→3)GlcNAc-, or the sialyl Le^x, NeuNAcα2→3Galß1→3 (Fucα1→4)GlcNAc-, determinants (Fukuda, <u>Biochim. Biophys. Acta</u> 780:119-150 (1985)). Such modifications are significant because these determinants, which are present on neutrophils and monocytes, serve as ligands for E- and P-selectin present on endothelial cells and platelets, respectively (see, for example, Larsen et al., <u>Cell</u> 63:467-474 (1990)).

In addition, tumor cells often express a significant amount of sialyl Le^x and/or sialyl Le^x on their cell surfaces. The interaction between E-selectin or P-selectin and these cell surface carbohydrates may play a role in tumor cell adhesion to endothelium during the metastatic process (Walz et al., supra, (1990)). Kojima et al., Biochem. Biophys. Res. Commun. 182:1288-1295 (1992) reported that selectin-dependent tumor cell adhesion to endothelial cells was abolished by blocking O-glycan synthesis. Complex sulfated O-glycans also may serve as ligands for the lymphocyte homing receptor, L-selectin

(Imai et al., J. Cell Biol. 113:1213-1221 (1991)).

These reported observations establish core 2 ß1→6 N-acetylglucosaminyltransferase as a critical enzyme in Oglycan biosynthesis. The availability of core 2 β 1 \rightarrow 6 N-5 acetylglucosaminyltransferase will allow the in vivo and in vitro production of specific glycoproteins having core 2 oligosaccharides and subsequent study of these variant Oglycans on cell-cell interactions. For example, core 2 B1→6 N-acetylglucosaminyltransferase is a useful marker for 10 transformed or cancerous cells. An understanding of the role of core 2 ßl→6 N-acetylglucosaminyltransferase in transformed and cancerous cells may elucidate a mechanism for the aberrant cell-cell interactions observed in these cells. In order to understand the control of expression of 15 these oligosaccharides and their function, isolation of a cDNA clone for core 2 \$1 -6 N-acetylglucosaminyltransferase is a prerequisite. However, the DNA sequence encoding core 2 ß1→6 N-acetylglucosaminyltransferase has not yet been reported.

Thus, a need exists for identifying the core 2 $\mathfrak{S}1\rightarrow 6$ N-acetylglucosaminyltransferase and the DNA sequences encoding this enzyme. The present invention satisfies this need and provides related advantages as well.

SUMMARY OF THE INVENTION

25 The present invention generally relates to a novel purified human $\mathfrak{S}1 \rightarrow 6$ N-acetylglucosaminyltransferase. A cDNA sequence encoding a 428 amino acid protein having B1→6 N-acetylglucosaminyltransferase activity also provided. The purified B1→6 human Nacetylglucosaminyltransferase, or an thereof, catalyzes the formation of critical branches in Oglycans.

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The invention further relates to a novel purified acceptor molecule, leukosialin, CD43, for core 2 ß1→6 N-acetylglucosaminyltransferase activity. The leukosialin cDNA encodes a novel variant leukosialin, which is created by alternative splicing of the genomic leukosialin DNA sequence.

Isolated nucleic acids encoding either core 2 \$1.76 N-acetylglucosaminyltransferase or leukosialin are disclosed, as are vectors containing the nucleic acids and recombinant host cells transformed with such vectors. The invention further provides methods of detecting such nucleic acids by contacting a sample with a nucleic acid probe having a nucleotide sequence capable of hybridizing with the isolated nucleic acids of the present invention.

The core 2 \$1.76 N-acetylglucosaminyltransferase and leukosialin amino acid and nucleic acid sequences disclosed herein can be purified from human cells or produced using well known methods of recombinant DNA technology.

The invention also discloses a method of isolating nucleic acid sequences encoding proteins that have an enzymatic activity. Such a nucleic acid sequence is obtained by transfecting the nucleic acid, which is contained within a vector having a polyoma virus replication origin, into a Chinese hamster ovary (CHO) cell line simultaneously expressing polyoma virus large T antigen and the acceptor molecule for the protein having an enzymatic activity.

BRIEF DESCRIPTION OF THE DRAWINGS

Figure 1 depicts the structures and biosynthesis of O-glycans. Structures of O-glycan cores can be classified into 4 groups (core 1 to core 4), each of which is synthesized starting with GalNAcαl→Ser/Thr. The core 1 structure is synthesized by the addition of a β1→3 Gal

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residue to the GalNAc residue. The core 1 structure can be converted to core 2 by the addition of a B1+6 Nacetylglucosaminyl residue. This intermediate is usually converted to the hexasaccharide by sequential addition of 5 galactose and sialic acid residues (bottom right). core 2 Bl-6 N-acetylglucosaminyltransferase and the limkage formed by the enzyme are indicated by a box. cell types, the core 2 structure can be extended by the addition of N-acetyllactosamine (Galßl→4GlcNAcßl→3) repeats 10 to form poly-N-acetyllactosamine. In the absence of core 2 B1→6 N-acetylglucosaminyltransferase, core 1 is converted to the monosial oform, then to the disial oform by sequential addition of $\alpha 2\rightarrow 3-$ and $\alpha 2\rightarrow 6-$ linked sialic acid residues (bottom left). Alternatively, core 3 can be synthesized by the addition of a Bl→3 N-acetylqlucosaminyl residue to the GalNAc residue. Core 3 can be converted to core 4 by another ß1→6 N-acetylglucosaminyltransferase figure).

Figure 2 depicts genomic DNA sequence (SEQ. ID. NO. 1) and cDNA sequence (SEQ. ID. NO. 2) of leukosialin. The genomic sequence is numbered relative to the transcriptional start site. Exon 1 and exon 2 have been previously described. Exon 1' is newly identified here. In the isolated cDNA, exon 1' is immediately followed by the exon 2 sequence. Deduced amino acids are presented under the coding sequence, which begins in exon 2 (SEQ. ID. NO. 3). A portion of the exon 2 sequence is shown.

Figure 3 establishes the ability of pGT/hCG to replicate in CHO cell lines expressing polyoma large T antigen and leukosialin. In panel A, six clonal CHO cell lines were examined for replication of pcDNAI-based pGT/hCG (lanes 1-6). In panel B, replication of cell clone 5 (CHO-Py-leu), was further examined by treatment with increasing concentrations of DpnI and XhoI (lanes 2 and 3). Plasmid DNA isolated from MOP-8 cells was used as a control (lane

1). Plasmid DNA was extracted using the Hirt procedure and samples were digested with XhoI and DpnI. In parallel, pGT/hCG plasmid purified from E. coli MC1061/P3 was digested with XhoI and DpnI (lane 7 in panel A and lane 4 in panel B) or XhoI alone (lane 8 in panel A and lane 5 in panel B). The arrow indicates the migration of plasmid DNA resistant to DpnI digestion. The arrowheads indicate plasmid DNA digested by DpnI.

Figure 4 shows the expression of T305 antigen expressed by pcDNAI-C2GnT. Subconfluent CHO-Py-leu cells were transfected with pcDNAI-C2GnT (panels A and B) or mock-transfected with pcDNAI (panels C and D). Sixty four hours after transfection, the cells were fixed, then incubated with mouse T305 monoclonal antibody followed by fluorescein isocyanate-conjugated sheep anti-mouse IgG (panels A, B and C). Two different areas are shown in panels A and B. Panel D shows a phase micrograph of the same field shown in panel C. Bar = 20µm.

Figure 5 depicts the cDNA sequence (SEQ. ID. NO. 20 4) and translated amino acid sequences (SEQ. ID. NO. 5) of core 2 \$\text{Sl} \to 6 \text{N-acetylglucosaminyltransferase}\$. The open reading frame and full-length nucleotide sequence of C2GnT are shown. The signal/membrane-anchoring domain is doubly underlined. The polyadenylation signal is boxed. 25 Potential \$N\$-glycosylation sites are marked with asterisks. The sequences are numbered relative to the translation start site.

Figure 6 shows the expression of core 2 β1+6 N-acetylglucosaminyltransferase mRNA in various cell types.

30 Poly(A)+ RNA (11 μg) from CHO-Py-leu cells (lane 1), HL-60 promyelocytes (lane 2), K562 erythrocytic cells (lane 3), and SP and L4 colonic carcinoma cells (lanes 4 and 5) was resolved by electrophoresis. RNA was transferred to a nylon membrane and hybridized with a radiolabeled fragment

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of pPROTA-C2GnT. Migration of RNA size markers is indicated.

Figure 7 illustrates the construction of the vector encoding the protein A-C2GnT fusion protein. The cDNA sequence corresponding to Pro³⁸ to His⁴²⁸ was fused in frame with the IgG binding domain of S. aureus protein A (bottom; SEQ. ID. NO. 6). The sequence includes the cleavable signal peptide, which allows secretion of the fused protein. The coding sequence is under control of the SV40 promoter. The remainder of the vector sequence shown was derived from rabbit ß-globin gene sequences, including an intervening sequence (IVS) and a polyadenylation signal (An).

DETAILED DESCRIPTION OF THE INVENTION

The present invention generally relates to a novel human core 2 β1+6 N-acetylglucosaminyltransferase. The invention further relates to a novel method of transient expression cloning in CHO cells that was used to isolate the cDNA sequence encoding human core 2 β1+6 N-acetylglucosaminyltransferase (C2GnT). The invention also relates to a novel human leukosialin, which is an acceptor molecule for core 2 β1+6 N-acetylglucosaminyltransferase activity.

Cells generally contain extremely low amounts of glycosyltransferases. As a result, cDNA cloning based on screening using an antibody or a probe based on the glycosyltransferase amino acid sequence has met with limited success. However, isolation of cDNAs encoding various glycosyltransferases can be achieved by transient expression of cDNA in recipient cells.

Successful application of the transient expression cloning method to isolate a cDNA sequence

encoding a glycosyltransferase requires an appropriate recipient cell line. Ideal recipient cells should not express the glycosyltransferase of interest. As a result, the recipient cells would normally lack the oligosaccharide structure formed by such a glycosyltransferase.

in the recipient cell line should result in formation of the specific oligosaccharide structure. The resultant oligosaccharide can be identified using a specific antibody or lectin that recognizes the structure. The recipient cell line also must support replication of an appropriate plasmid vector.

COS-1 cells initially appear to satisfy the requirements for using the transient expression method. 15 COS-1 cells express SV40 large T antigen and support the replication of plasmid vectors harboring a SV40 replication origin (Gluzman et al., Cell 23:175-182 (1981)). Although COS-1 cells, themselves, express a glycosyltransferases, COS-1 cells have been used to clone cDNA sequences encoding human blood group Lewis $\alpha 1 \rightarrow 3/4$ fucosyltransferase and murine α1→3 galactosyltransferase (Kukowska-Latallo et al., Genes and Devel. 4:1288-1303 (1990); Larsen et al., Proc. Natl. Acad. Sci. USA 86:8227-8231 (1989)). Also, Goelz et al., Cell 63:175-182 (1990), utilized an antibody that inhibits E-selectin mediated adhesion to isolate a cDNA sequence encoding $\alpha 1\rightarrow 3$ fucosyltransferase.

An attempt was made to use COS-1 cells to isolate CDNA clones encoding B1→6 core 2 N-30 acetylglucosaminyltransferase. cos-1 cells were transfected using cDNA obtained from activated human T cells, which 2 express the core B1→6 Nacetylglucosaminyltransferase. Transfected cells suspected of expressing core 2 \$1.46 N-acetylglucosaminyltransferase WO 94/07917 PCT/US93/093/B

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in the transfected cells were identified by the presence of increased levels of the core 2 oligosaccharide structure formed by core 2 Bl→6 N-acetylglucosaminyltransferase activity. The presence of the core 2 structure was identified using the monoclonal antibody, T305, which identifies a hexasaccharide on leukosialin. A clone expressing high levels of the T305 antigen was isolated and sequenced.

Surprisingly, transfection using COS-1 cells resulted in the isolation of a cDNA clone encoding a nowel variant of human leukosialin, which is the acceptor molecule for core 2 Bl-6 N-acetylglucosaminyltransferase activity. Examination of the cDNA sequence of the newly isolated leukosialin revealed the cDNA sequence was formed as a result of alternative splicing of exons in the genomic leukosialin DNA sequence. Specifically, the newly isolated leukosialin is encoded by cDNA sequence containing a previously undescribed non-coding exon at the 5'-terminus (exon 1' in Figure 2; SEQ. ID. NO. 1 and SEQ. ID. NO. 2).

20 The unexpected result obtained using COS-1 cells led to the development of a new transfection system to isolate a cDNA sequence encoding core 2 ß1→6 N-acetylglucosaminyltransferase. CHO cells, which do not normally express the T305 antigen, were transfected with DNA sequences encoding human leukosialin and the polyoma virus large T antigen. A cell line, designated CHO-Py-leu, which expresses human leukosialin and polyoma virus large T antigen, was isolated.

CHO-Py-leu cells were used for transient 30 expression cloning of a cDNA sequence encoding core 2 ₺1→6 N-acetylglucosaminyltransferase. CHO-Py-leu cells were transfected with CDNA obtained from human HL-60 promyelocytes. A plasmid, pcDNAI-C2Gnt, which directed expression of the T305 antigen, was isolated and the cDNA insert was sequenced (see Figure 5; SEQ. ID. NO. 4). The 2105 base pair cDNA sequence encodes a putative 428 amino acid protein. The genomic DNA sequence encoding can be isolated using methods well known to those skilled in the art, such as nucleic acid hybridization using the core 2 B1+6 N-acetylglucosaminyltransferase cDNA disclosed herein to screen, for example, a genomic library prepared from HL-60 promyelocytes.

An enzyme similar to the disclosed human core 2 10 Bl→6 N-acetylglucosaminyltransferase has been purified from bovine tracheal epithelium (Ropp et al., J. Biol. Chem. 266:23863-23871 (1991), which is incorporated herein by reference. The apparent molecular weight of the bovine enzyme is ~69kDa. In comparison, the predicted molecular 15 weight of the polypeptide portion of core 2 β 1 \rightarrow 6 Nacetylglucosaminyltransferase is ~50kDa. The deduced amino acid sequence o f 2 B 1 → 6 core N acetylglucosaminyltransferase reveals two to potential N-glycosylation sites, suggesting N-glycosylation 20 and O-glycosylation, other or post-translational modification, could account for the larger apparent size of the bovine enzyme.

Expression of the cloned C2GnT sequence, or a fragment thereof, directed formation of the specific O-glycan core 2 oligosaccharide structure. Although several cDNA sequences encoding glycosyltransferases have been isolated (Paulson and Colley, J. Biol. Chem. 264:17615-17618 (1989); Schachter, Curr. Opin. Struct. Biol. 1:755-765 (1991), which are incorporated herein by reference), C2GnT is the first reported cDNA sequence encoding an enzyme involved exclusively in O-glycan synthesis.

In O-glycans, $\beta 1 \rightarrow 6$ N-acetylglucosaminyl linkages may occur in both core 2, $Gal\beta 1 \rightarrow 3(GlcNAc\beta 1 \rightarrow 6)GalNAc$, and core 4, $GlcNAc\beta 1 \rightarrow 3(GlcNAc\beta 1 \rightarrow 6)GalNAc$, structures

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(Brockhausen et al., <u>Biochemistry</u> 24:1866-1874 (1985), which is incorporated herein by reference. In addition, β1→6 N-acetylglucosaminyl linkages occur in the side chains of poly-N-acetyllactosamine, forming the I-structure (Piller et al., <u>J. Biol. Chem.</u> 259:13385-13390 (1984), which is incorporated herein by reference), and in the side chain attached to α-mannose of the N-glycan core structure, forming a tetraantennary saccharide (Cummings et al., <u>J. Biol. Chem.</u> 257:13421-13427 (1982), which is incorporated herein by reference). The enzymes responsible for these linkages all share the unique property that Mn²+ is not required for their activity.

Although it was originally suggested that these \$1+6 N-acetylglucosaminyl linkages were formed by the same enzyme (Piller at al., 1984), the present disclosure clearly demonstrates that the HL-60-derived core 2 \$1+6 N-acetylglucosaminyltransferase is specific for the formation only of O-glycan core 2. This result is consistent with a recent report demonstrating that myeloid cell lysates contain the enzymatic activity associated with core 2, but not core 4, formation (Brockhausen et al., supra, (1991)).

Analysis of mRNA isolated from colonic cancer cells indicated core 2 Bl > 6 N-acetylglucosaminyltransferase is expressed in these cells. Recent studies using affinity 25 absorption suggested at least two different $\mathfrak{S}1 \rightarrow 6$ Nacetylglucosaminyltransferases were present in tracheal epithelium (Ropp et al., supra, (1991)). One of these transferases formed core 2, core 4, and I structures. Thus, at least one other B 1 → 6 acetylglucosaminyltransferase present in epithelial cells 30 can form core 2, core 4 and I structures. Similarly, a B1→6 N-acetylglucosaminyltransferase present in Novikoff hepatoma cells can form both core 2 and I structures (Koenderman et al., Eur. J. Biochem. 166:199-208 (1987), 35 which is incorporated herein by reference).

The acceptor molecule specificity of core 2 ß1→6 N-acetylglucosaminyltransferase is different from the specificity of the enzymes present in tracheal epithelium Thus, a family of $\beta 1 \rightarrow 6$ Nand Novikoff hepatoma cells. 5 acetylglucosaminyltransferases can exist, the members of which differ in acceptor specificity but are capable of forming the same linkage. Members of this family are isolated from cells expressing B1 → 6 acetylqlucosaminyltransferase activity, using, for example, 10 nucleic acid hybridization assays and studies of acceptor molecule specificity. Such a family was reported for the αl→3 fucosyltransferases (Weston et al., J. Biol. Chem. 267:4152-4160 (1992), which is incorporated herein by reference).

15 The formation of the core 2 structure is critical to cell structure and function. For example, the core 2 structure is essential elongation of for acetyllactosamine and for formation of sialyl Lex or sialyl Le' structures. Furthermore, the biosynthesis of cartilage 20 keratan sulfate may be initiated by the core 2 β 1 \rightarrow 6 Nacetylglucosaminyltransferase, since the keratan sulfate chain is extended from a branch present in core 2 structure in the same way as poly-N-acetyllactosamine (Dickenson et al., Biochem. J. 269:55-59 (1990), which is incorporated 25 herein by reference). Keratan sulfate is absent in wildtype CHO cells, which do not express the core 2 β 1 \rightarrow 6 Nacetylglucosaminyltransferase (Esko et al., J. Biol. Chem. 261:15725-15733 (1986), which is incorporated herein by reference). These structures are believed to be important 30 for cellular recognition and matrix formation. availability of the cDNA clone encoding the core 2 β 1 \rightarrow 6 Nacetylglucosaminyltransferase will aid in understanding how the various carbohydrate structures are formed during differentiation and malignancy. Manipulation of expression of the various carbohydrate structures by gene

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transfer and gene inactivation methods will help elucidate the various functions of these structures.

The present invention is directed to a method for transient expression cloning in CHO cells of cDNA sequences encoding proteins having enzymatic activity. Isolation of human core 2 \$\text{Sl} \to 6 \text{N-acetylglucosaminyltransferase} is provided as an example of the disclosed method. However, the method can be used to obtain cDNA sequences encoding other proteins having enzymatic activity.

10 For example, lectins and antibodies reactive with other specific oligosaccharide structures are available and can be used to screen for glycosyltransferase activity. Also, CHO cell lines that have defects in glycosylation have been isolated. These cell lines can be used to study 15 the activity of the corresponding glycosyltransferase (Stanley, Ann. Rev. Genet. 18:525-552 (1984), which is incorporated herein by reference). CHO cell lines also have been selected for various defects in cellular metabolism, loss of expression of cell surface molecules 20 and resistance to cytotoxic drugs (see, for example, Malmström and Krieger, J. Biol. Chem. 266:24025-24030 (1991); Yayon et al., Cell 64:841-848 (1991), which are incorporated herein by reference). The approach disclosed herein should allow isolation of cDNA sequences encoding 25 the proteins involved in these various cellular functions.

As used herein, the terms "purified" and "isolated" mean that the molecule or compound is substantially free of contaminants normally associated with a native or natural environment. For example, a purified protein can be obtained from a number of methods. The naturally-occurring protein can be purified by any means known in the art, including, for example, by affinity purification with antibodies having specific reactivity with the protein. In this regard, anti-core 2 \$1.000 N-

acetylglucosaminyltransferase antibodies can be used to substantially purify naturally-occurring core 2 $\mathfrak{S}1\rightarrow 6$ N-acetylglucosaminyltransferase from human HL-60 promyelocytes.

Alternatively, a purified protein of the present invention can be obtained by well known recombinant methods, utilizing the nucleic acids disclosed herein, as described, for example, in Sambrook et al., Molecular Cloning: A Laboratory Manual 2d ed. (Cold Spring Harbor Laboratory 1989), which is incorporated herein by reference, and by the methods described in the Examples below. Furthermore, purified proteins can be synthesized by methods well known in the art.

As used herein, the phrase "substantially the sequence" includes the described nucleotide or amino acid 15 sequence and sequences having one or more additions, deletions or substitutions that do not substantially affect the ability of the sequence to encode a protein have a desired functional activity. In addition, the phrase 20 encompasses any additional sequence that hybridizes to the disclosed sequence under stringent hybridization sequences. Methods of hybridization are well known to those skilled in the art. For example, sequence modifications that do not substantially alter such activity are intended. 25 protein having substantially the amino acid sequence of Figure 5 (SEQ. ID. NO. 5) refers to core 2 β 1+6 Nacetylglucosaminyltransferase encoded by the cDNA described in Example IV, as well as proteins having amino acid sequences that are modified but, nevertheless, retain the 30 functions of core 2 B1→6 N-acetylqlucosaminyltransferase. One skilled in the art can readily determine such retention of function following the quidance set forth, for example, in Examples V and VI.

The present invention is further directed to active fragments of the human core acetylglucosaminyltransferase protein. As used herein, an active fragment refers to portions of the protein that 5 substantially retain the glycosyltransferase activity of the intact core 2 B1→6 N-acetylglucosaminyltransferase One skilled in the art can readily identify protein. active fragments of proteins such as core 2 ß1→6 Nacetylglucosaminyltransferase by comparing the activities 10 of a selected fragment with the intact protein following the guidance set forth in the Examples below.

As used herein, the term "glycosyltransferase activity" refers to the function of a glycosyltransferase to link sugar residues together through a glycosidic bond 15 to oligosaccharides. create critical branches in Glycosyltransferase activity results in the transfer of a monosaccharide to an appropriate acceptor molecule, such that the acceptor molecule contains oligosaccharides having critical branches. One skilled in 20 the art would understand the terms "enzymatic activity" and "catalytic activity" to generally refer to a function of certain proteins, such as the function of those proteins having glycosyltransferase activity.

As used herein, the term "acceptor molecule" 25 refers to a molecule that is acted upon by a protein having enzymatic activity. For example, an acceptor molecule, such as leukosialin, as identified by the amino acid sequence of Figure 2 (SEQ. ID. NO. 3), accepts the transfer of a monosaccharide due to glycosyltransferase activity. 30 An acceptor molecule, such as leukosialin, may already The transfer of contain one or more sugar residues. monosaccharides molecule, to an acceptor leukosialin, results in the formation of critical branches of oligosaccharides.

As used herein, the term "critical branches" refers to oligosaccharide structures formed by specific glycosyltransferase activity. Critical branches may be involved in various cellular functions, such as cell-cell recognition. The oligosaccharide structure of a critical branch can be determined using methods well known in the art, such as the method for determining the core 2 oligosaccharide structure, as described in Examples V and VI.

Relatedly, the invention also provides nucleic 10 B1→6 2 acids encoding the human core leukosialin and acetylglucosaminyltransferase protein protein described above. The nucleic acids can be in the form of DNA, RNA or cDNA, such as the novel C2GnT cDNA of 15 2105 base pairs identified in Figure 5 (SEQ. ID. NO. 4) or the novel leukosialin cDNA identified in Figure 2 (SEQ. ID. Such nucleic acids can also be NO. 2), for example. chemically synthesized by methods known in the art, including, for example, the use of an automated nucleic 20 acid synthesizer.

The nucleic acid can have substantially the nucleotide sequence of C2GnT, identified in Figure 5 (SEQ. ID. NO. 4), or leukosialin identified in Figure 2 (SEQ. ID. NO. 2). Portions of such nucleic acids that encode active fragments of the core 2 \$1 \to 6 N - acetylglucosaminyltransferase protein or leukosialin protein of the present invention also are contemplated.

Nucleic acid probes capable of hybridizing to the nucleic acids of the present invention under reasonably stringent conditions can be prepared from the cloned sequences or by synthesizing oligonucleotides by methods known in the art. The probes can be labeled with markers according to methods known in the art and used to detect the nucleic acids of the present invention. Methods for

detecting such nucleic acids can be accomplished by contacting the probe with a sample containing or suspected of containing the nucleic acid under hybridizing conditions, and detecting the hybridization of the probe to the nucleic acid.

The present invention is further directed to vectors containing the nucleic acids described above. term "vector" includes vectors that are capable expressing nucleic acid sequences operably linked to 10 regulatory sequences capable of effecting their expression. Numerous cloning vectors are known in the art. selection of an appropriate cloning vector is a matter of choice. In general, useful vectors for recombinant DNA are often plasmids, which refer to circular double stranded DNA 15 loops such as pcDNAI or pcDSRα. As used herein, "plasmid" and "vector" may be used interchangeably as the plasmid is a common form of a vector. However, the invention is intended to include other forms of expression vectors that serve equivalent functions.

20 Suitable host cells containing the vectors of the present invention are also provided. Host cells can be transformed with a vector and used to express the desired recombinant or fusion protein. Methods of recombinant expression in a variety of host cells, such as mammalian, 25 yeast, insect or bacterial cells are widely known. For example, a nucleic acid encoding core 2 ß1→6 acetylglucosaminyltransferase or a nucleic acid encoding leukosialin can be transfected into cells using the calcium phosphate technique or other transfection methods, such as 30 those described in Sambrook et al., supra, (1989).

Alternatively, nucleic acids can be introduced into cells by infection with a retrovirus carrying the gene or genes of interest. For example, the gene can be cloned into a plasmid containing retroviral long terminal repeat

sequences, the C2Gnt DNA sequence or the leukosialin DNA sequence, and an antibiotic resistance gene for selection. The construct can then be transfected into a suitable cell line, such as PA12, which carries a packaging deficient provirus and expresses the necessary components for virus production, including synthesis of amphotrophic glycoproteins. The supernatant from these cells contain infectious virus, which can be used to infect the cells of interest.

10 Isolated recombinant polypeptides or proteins can be obtained by growing the described host cells under conditions that favor transcription and translation of the transfected nucleic acid. Recombinant proteins produced by the transfected host cells are isolated using methods set forth herein and by methods well known to those skilled in the art.

Also provided are antibodies having specific N-B1 → 6 2 core the with reactivity leukosialin acetylglucosaminyltransferase protein or 20 protein of the present invention. Active fragments of antibodies, for example, Fab and Fab'2 fragments, having specific reactivity with such proteins are intended to fall within the definition of an "antibody." Antibodies exhibiting a titer of at least about 1.5×10^5 , as 25 determined by ELISA, are useful in the present invention.

The antibodies of the invention can be produced by any method known in the art. For example, polyclonal and monoclonal antibodies can be produced by methods described in Harlow and Lane, Antibodies: A Laboratory

Manual (Cold Spring Harbor 1988), which is incorporated herein by reference. The proteins, particularly core 2

B1+6 N-acetylglucosaminyltransferase or leukosialin of the present invention can be used as immunogens to generate such antibodies. Altered antibodies, such as chimeric,

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humanized, CDR-grafted or bifunctional antibodies can also be produced by methods well known to those skilled in the art. Such antibodies can also be produced by hybridoma, chemical synthesis or recombinant methods described, for example, in Sambrook et al., <u>supra</u>, (1989).

The antibodies can be used for determining the presence or purification of the core 2 \$1→6 N-acetylglucosaminyltransferase protein or the leukosialin protein of the present invention. With respect to the detecting of such proteins, the antibodies can be used for in vitro or in vivo methods well known to those skilled in the art.

Finally, kits useful for carrying out the methods of the invention are also provided. The kits can contain a core 2 \$\text{Bl} \to 6 N\$-acetylglucosaminyltransferase protein, antibody or nucleic acid of the present invention and an ancillary reagent. Alternatively, the kit can contain a leukosialin protein, antibody or nucleic acid of the present invention and an ancillary reagent. An ancillary reagent may include diagnostic agents, signal detection systems, buffers, stabilizers, pharmaceutically acceptable carriers or other reagents and materials conventionally included in such kits.

A cDNA sequence encoding core 2 \$1 \to 6 N-25 acetylglucosaminyltransferase was isolated and core 2 \$1 \to 6 N-acetylglucosaminyltransferase activity was determined. This is the first report of transient expression cloning using CHO cells expressing polyoma large T antigen. The following examples are intended to illustrate but not limit the present invention.

EXAMPLE I

EXPRESSION CLONING IN COS-1 CELLS OF THE cDNA FOR THE PROTEIN CARRYING THE HEXASACCHARIDES

COS-1 cells were transfected with a cDNA library, 5 pcDSRα-2F1, constructed from poly(A)+ RNA of activated T B1→6 lymphocytes, which express the core 2 acetylglucosaminyltransferase (Yokota et al., Proc. Natl. Acad. Sci. USA 83:5894-5898 (1986); Piller et al., supra, (1988), which are incorporated herein by reference). COS-1 10 cells support replication of the pcDSRa constructs, which contain the SV40 replication origin. Transfected cells were selected by panning using monoclonal antibody T305, which recognizes sialylated branched hexasaccharides (Piller et al., supra, (1991); Saitoh et al., supra, 15 (1991)). Methods referred to in this example are described in greater detail in the examples that follow.

Following several rounds of transfection, one plasmid, pcDSRα-leu, directing high expression of the T305 antigen was identified. The cloned cDNA insert was isolated and sequenced, then compared with other reported sequences. The newly isolated cDNA sequence was nearly identical to the sequence reported for leukosialin, except the 5'-flanking sequences were different (Pallant et al., Proc. Natl. Acad. Sci. USA 86:1328-1332 (1989), which is incorporated herein by reference).

Comparison of the cloned cDNA sequence with the genomic leukosialin DNA sequence revealed the start site of the cDNA sequence is located 259 bp upstream of the transcription start site of the previously reported sequence (Figure 2; compare Exon 1' and Exon 1) (Shelley et al., Biochem. J. 270:569-576 (1990); Kudo and Fukuda, J. Biol. Chem. 266:8483-8489 (1991), which are incorporated herein by reference). A consensus splice site was

identified at the exon-intron junction of the newly identified 122 bp exon 1' in pcDSRα-leu (Breathnach and Chambon, Ann. Rev. Biochem. 50:349-383 (1981), which is incorporated herein by reference). This splice site is followed by the exon 2 sequence.

results indicate the T305 antibody preferentially binds to branched hexasaccharides attached to leukosialin. Indeed, a small amount of the hexasaccharides (approximately 8% of the total) was 10 detected in O-glycans isolated from control COS-1 cells. T305 binding is similar to anti-M and anti-N antibodies, which recognize both the glycan and polypeptide portions of erythrocyte glycoprotein, glycophorin (Sadler et al., J. Biol. Chem 254: 2112-2119 (1979), which is incorporated 15 herein by reference). These observations are consistent with reports that only leukosialin strongly reacted with T305 in Western blots of leukocyte cell extracts, even though leukocytes also express other glycoproteins, such as CD45, that must also contain the same hexasaccharides 20 (Piller et al., supra, (1991); Saitoh et al., supra, (1991)).

EXAMPLE II

ESTABLISHMENT OF CHO CELL LINES THAT STABLY EXPRESS POLYOMA VIRUS LARGE T ANTIGEN AND LEUKOSIALIN

25 T305 preferentially binds to branched hexasaccharides attached to leukosialin. Such hexasaccharides are not present on the erythropoietin glycoprotein produced in CHO cells, although the glycoprotein does contain the precursor tetrasaccharide 30 (Sasaki et al., <u>J. Biol. Chem.</u> 262:12059-12076 (1987), which is incorporated herein by reference). T305 antigen also is not detectable in CHO cells transiently transfected with $pcDSR\alpha$ -leu. In order to screen for the presence of a cDNA clone expressing core 2 \$1→6 N-acetylglucosaminyltransferase activity, a CHO cell line expressing both leukosialin and polyoma large T antigen was established (see, for example, Heffernan and Dennis Nucl. Acids Res. 19:85-92 (1991), which is incorporated herein by reference).

A plasmid vector, pPSVE1-PyE, which contains Vectors: . the polyoma virus early genes under the control of the SV40 early promoter, was constructed using a modification of the 10 method of Muller et al., Mol. Cell. Biol. 4:2406-2412 (1984), which is incorporated herein by reference. Plasmid pPSVE1 was prepared using pPSG4 (American Type Culture Collection 37337) and SV40 viral DNA (Bethesda Research laboratories) essentially as described by Featherstone et al., Nucl. Acids Res. 12:7235-7249 (1984), which incorporated herein by reference. Following EcoRI and HincII digestion of plasmid pPyLT-1 (American Type Culture Collection 41043), a DNA sequence containing the carboxy terminal coding region of polyoma virus large T antigen was 20 isolated. The HincII site was converted to an EcoRI site by blunt-end ligation of phosphorylated EcoRI linkers Plasmid pPSVE1-PyE was generated by (Stratagene). inserting the carboxy-terminal coding sequence for large T antigen into the unique EcoRI site of plasmid pPSVE1.

constructed by pZIPNEO-leu was 25 Plasmid introducing the EcoRI fragment of PEER-3 cDNA, which coding sequence complete contains the leukosialin, into the unique EcoRI site of plasmid pZIPNEO (1984), which is (Cepko et al., <u>Cell</u> 37:1053-1063 30 incorporated herein by reference). Plasmid structures were confirmed by restriction mapping and by sequencing the construction sites. pZIPNEO was kindly provided by Dr. Channing Der.

Transfection: CHODG44 cells were grown in 100 mm tissue

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culture plates. When the cells were 20% confluent, they were co-transfected with a 1:4 molar ratio of pZIPNEO-leu and pPSVE1-PyE using the calcium phosphate technique (Graham and van der Eb, <u>Virology</u> 52:456-467 (1973), which is incorporated herein by reference). Transfected cells were isolated and maintained in medium containing 400 µg/ml G-418 (active drug).

Leukosialin expression:

The total pool of G418resistant transfectants was enriched for human leukosialin
expressing cells by a one-step panning procedure using
anti-leukosialin antibodies and goat anti-rabbit IgG coated
panning dishes (Sigma) (Carlsson and Fukuda J. Biol. Chem.
261:12779-12786 (1986), which is incorporated herein by
reference). Clonal cell lines were obtained by limiting
dilution. Six clonal cell lines expressing human
leukosialin on the cell surface were identified by indirect
immunofluorescence and isolated for further studies
(Williams and Fukuda J. Cell Biol. 111:955-966 (1990),
which is incorporated herein by reference).

Polyoma virus-mediated replication: The ability of the six clonal cell lines to support polyoma virus large T antigen-mediated replication of plasmids was assessed by determining the methylation status of transfected plasmids containing a polyoma virus origin of replication (Muller at al., supra, 1984; Heffernan and Dennis, supra, 1991). Plasmid pGT/hCG contains a fused β1+4 galactosyltransferase and human chorionic gonadotropin α-chain DNA sequence inserted in plasmid pcDNAI, which contains a polyoma virus replication origin (Aoki et al., Proc. Natl. Acad. Sci., USA 89, 4319-4323 (1992), which is incorporated herein by reference).

Plasmid pGT/hCG was isolated from methylase-positive $E.\ coli$ strain MC1061/P3 (Invitrogen), which methylates the adenine residues in the DpnI recognition

site, "GATC". The methylated DpnI recognition site is susceptible to cleavage by DpnI. In contrast, the DpnI recognition site of plasmids replicated in mammalian cells is not methylated and, therefore, is resistant to DpnI digestion.

Methylated plasmid pGT/hCG was transfected by lipofection into each of the six selected clonal cell lines expressing leukosialin. After 64 hr, low molecular weight plasmid DNA was isolated from the cells using the method of Hirt, J. Mol. Biol. 26:365-369 (1967), which is incorporated herein by reference. Isolated plasmid DNA was digested with XhoI and DpnI (Stratagene), subjected to electrophoresis in a 1% agarose gel, and transferred to nylon membranes (Micron Separations Inc., MA).

the B1→4 fragment of 15 Α 0.4 kbSmaI galactosyltransferase sequence of pGT/hCG was DNA radiolabeled with [32P]dCTP using the random primer method (Feinberg and Vogelstein, Anal. Biochem. 132:6-13 (1983), which is incorporated herein by reference). Hybridization 20 was performed using methods well-known to those skilled in the art (see, for example, Sambrook et al., supra, (1989)). Following hybridization, the membranes were washed several times, including a final high stringency wash in 0.1 x SSPE, 0.1% SDS for 1 hr at 65°C, then exposed to Kodak X-AR 25 film at -70°C.

Four of the six clones tested supported replication of the pcDNAI-based plasmid, pGT/hCG (Fig. 3.A., lanes 1, 3, 4 and 5). MOP-8 cells, a 3T3 cell line transformed by polyoma virus early genes (Muller et al., supra, (1984)), expresses endogenous core 2 ß1→6 N-acetylglucosaminyltransferase activity and was used as a control for the replication assay (Fig. 3.B., lane 1). One clonal cell line that supported pGT/hCG replication, CHO-Py-leu (Fig. 3.A., lane 5; Fig. 3.B., lanes 2 and 3) and

expressed a significant amount of leukosialin, was selected for further studies. pGT/hCG was kindly provided by Dr. Michiko Fukuda.

EXAMPLE III

5 ISOLATION OF A cDNA SEQUENCE DIRECTING EXPRESSION OF THE HEXASACCHARIDE ON LEUKOSIALIN

Poly(A)⁺ RNA was isolated from HL-60 promyelocytes, which contain a significant amount of the core 2 \$1.000 acetylglucosaminyltransferase (Saitoh et al., supra, (1991)). A cDNA expression library, pcDNAI-HL-60, was prepared (Invitrogen) and the library was screened for clones directing the expression of the T305 antigen.

Plasmid DNA from the pcDNAI-HL-60 cDNA library was transfected into CHO-Py-leu cells using a modification of the lipofection procedure, described below (Felgner et al., Proc. Natl. Acad. Sci. USA 84:7413-7417 (1987), which is incorporated herein by reference). CHO-Py-leu cells were grown in 100 mm tissue culture plates. When the cells were 20% confluent, they were washed twice with Opti-MEM I (GIBCO). Fifty μg of lipofectin reagent (Bethesda Research Laboratories) and 20 μg of purified plasmid DNA were each diluted to 1.5 ml with Opti-MEM I, then mixed and added to the cells. After incubation for 6 hr at 37°C, the medium was removed, 10 ml of complete medium was added and incubation was continued for 16 hr at 37°C. The medium was then replaced with 10 ml of fresh medium.

Following a 64 hr period to allow transient expression of the transfected plasmids, the cells were detached in PBS/5mM EDTA, pH7.4, for 30 min at 37°C, pooled, centrifuged and resuspended in cold PBS/10mM

EDTA/5% fetal calf serum, pH7.4, containing a 1:200 dilution of ascites fluid containing T305 monoclonal antibody. The cells were incubated on ice for 1 hr, then washed in the same buffer and panned on dishes coated with goat anti-mouse IgG (Sigma) (Wysocki and Sato Proc. Natl. Acad. Sci. USA 75:2844-2848 (1978); Seed & Aruffo Proc. Natl. Acad. Sci. USA 84:3365-3369 (1987), which are incorporated herein by reference). T305 monoclonal antibody was kindly provided by Dr. R.I. Fox, Scripps Research Foundation, La Jolla, CA.

Plasmid DNA was recovered from adherent cells by the method of Hirt, supra, (1967), treated with DpnI to eliminate plasmids that had not replicated in transfected cells, and transformed into E. coli strain MC1061/P3.

Plasmid DNA was then recovered and subjected to a second round of screening. E. coli transformants containing plasmids recovered from this second enrichment were plated to yield 8 pools of approximately 500 colonies each. Replica plates were prepared using methods well-known to those skilled in the art (see, for example, Sambrook et al., supra, (1989)).

The pooled plasmid DNA was prepared from replica plates and transfected into CHO-Py-leu cells. The transfectants were screened by panning. One plasmid pool was selected and subjected to three subsequent rounds of selection. One plasmid, pcDNAI-C2GnT, which directed the expression of the T305 antigen, was isolated. CHO-Py-leu cells transfected with pcDNAI-C2GnT express the antigen recognized by T305, whereas CHO-Py-leu cells transfected with pcDNAI are negative for T305 antigen (Fig. 4). These results show pcDNAI-C2GnT directs the expression of a new determinant on leukosialin that is recognized by T305 monoclonal antibody. This determinant is the branched h e x a s a c c h a r i d e s e q u e n c e , NeuNAcc2→3Galßl→3(NeuNAcc2→3Galßl→4 GlcNAcßl→6)GalNAc.

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EXAMPLE IV

CHARACTERIZATION OF C2GnT

DNA sequence: The cDNA insert in plasmid pcDNAI-C2GnT was sequenced by the dideoxy chain termination method using 5 Sequenase version 2 reagents (United States Biochemicals) (Sanger et al., Proc. Natl. Acad. Sci. USA 74:5463-5467 (1977), which is incorporated herein by reference). Both strands were sequenced using 17-mer synthetic oligonucleotides, which were synthesized as the sequence of the cDNA insert became known.

Plasmid pcDNAI-C2GnT contains a 2105 base pair insert (Fig. 5). The cDNA sequence ends 1878 bp downstream of the putative translation start site. A polyadenylation signal is present at nucleotides 1694-1699. The significance of the large number of nucleotides between the polyadenylation signal and the beginning of the polyadenyl chain is not clear. However, this sequence is A/T rich.

Deduced amino acid sequence: The cDNA insert in plasmid pcDNAI-C2GnT encodes a single open reading frame in the sense orientation with respect to the pcDNAI promoter (Fig. 5). The open reading frame encodes a putative 428 amino acid protein having a molecular mass of 49,790 daltons.

Pydropathy analysis indicates the predicted protein is a type II transmembrane molecule, as are all previously reported mammalian glycosyltransferases (Schachter, supra, (1991)). In this topology, a nine amino acid cytoplasmic NH2-terminal segment is followed by a 23 amino acid transmembrane domain flanked by basic amino acid residues. The large COOH-terminus consists of the stem and catalytic domains and presumably faces the lumen of the Golgi complex.

The putative protein contains three potential N-glycosylation sites (Fig. 5, asterisks). However, one of these sites contains a proline residue adjacent to asparagine and is not likely utilized <u>in vivo</u>.

No matches were obtained when the C2GnT cDNA sequence and deduced amino acid sequence were compared with sequences listed in the PC/Gene 6.6 data bank. In particular, no homology was revealed between the deduced amino acid sequence of C2GnT and other glycosyltransferases, including N-acetylglucosaminyltransferase I (Sarkar et al., Proc. Natl. Acad. Sci. USA 88:234-238 (1991), which is incorporated herein by reference).

mRNA expression: Poly(A) * RNA was prepared using a kit 15 (Stratagene) and resolved by electrophoresis on a 1.2% agarose/2.2 M formaldehyde gel, and transferred to nylon membranes (Micro Separations Inc., MA) using methods wellknown to those skilled in the art (see, for example, Sambrook et al., supra, (1989)). Membranes were probed 20 using the EcoRI insert of pPROTA-C2GnT (see below) radiolabeled with [32P]dCTP by the random priming method (Feinberg and Vogelstein, supra, (1983). Hybridization was performed in buffers containing 50% formamide for 24 hr at 42°C (Sambrook et al., <u>supra</u>, (1989)). Following 25 hybridization, filters were washed several times 1xSSPE/0.1% SDS at room temperature and once 0.1xSSPE/0.1% SDS at 42°C, then exposed to Kodak X-AR film at -70°C.

Fig. 6 compares the level of core 2 \$1.76 N-30 acetylglucosaminyltransferase mRNA isolated from HL-60 promyelocytes, K562 erythroleukemia cells, and poorly metastatic SP and highly metastatic L4 colonic carcinoma cells. The major RNA species migrates at a size essentially identical to the -2.1 kb C2GnT cDNA sequence.

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The same result is observed for HL-60 cells and the two colonic cell lines, which apparently synthesize the hexasaccharides. In addition, two transcripts of -3.3 kb and 5.4 kb in size were detected in these cell lines. The two larger transcripts may result from differential usage of polyadenylation signals.

No hybridization occurred with poly(A)+ RNA isolated from K562 cells, which lack the hexasaccharide, but synthesize the tetrasaccharide (Carlsson et al., supra, (1986)), which is incorporated herein by reference. Similarly, no hybridization was observed for poly(A)+ RNA isolated from CHO-Py-leu cells (Fig. 6, lane 1).

EXAMPLE V

EXPRESSION OF ENZYMATICALLY ACTIVE \$1+6 N-ACETYLGLUCOSAMINYLTRANSFERASE

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In order to confirm that C2GnT cDNA encodes for core 2 \$1.76 N-acetylglucosaminyltransferase, enzymatic activity was examined in CHO-Py-leu cells transfected with pcDNAI or pcDNAI-C2GnT. Following a 64 hr period to allow transient expression, cell lysates were prepared and core 2 \$1.76 N-acetylglucosaminyltransferase activity was measured.

N-acetylglucosaminyltransferase assays were performed essentially as described by Saitoh et al., supra, (1991), Yousefi et al., supra, (1991), and Lee et al., J. Biol. Chem. 265:20476-20487 (1990), which is incorporated herein by reference. Each reaction contained 50 mM MES, pH7.0, 0.5 μCi of UDP-[³H]GlcNAc in 1 mM UDP-GlcNAc, 0.1 M GlcNAc, 10 mM Na,EDTA, 1mM of acceptor and 25 μl of either cell lysate, cell supernatant or IgG-Sepharose matrix in a total reaction volume of 50 μl.

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Reactions were incubated for 1 hr at 37°C, then processed by C18 Sep-Pak chromatography (Waters) (Palcic et al., J. Biol. Chem. 265:6759-6769 (1990), which is incorporated herein by reference). Core 2 and core 4 £1.65 N-acetylglucosaminyltransferase were assayed using the acceptors p-nitrophenyl Gal£1.3GalNAc and p-nitrophenyl GlcNAc£1.3GalNAc, respectively (Toronto Research Chemicals).

UDP-GlcNAc: α-Man βl→6 N
acetylglucosaminyltransferase(V) was assayed using the acceptor GlcNAcβl→2Manαl→6Glc-β-O-(CH₂),CH₃. The blood group I enzyme, UDP-GlcNAc:GlcNAcβl→3Galβl→4GlcNAc (GlcNAc to Gal) βl→6 N-acetylglucosaminyltransferase, was assayed using GlcNAcβl→3Galβl→4GlcNAcβl→6Manβl→O-(CH₂),COOCH₃

or Galβl→4GlcNAcβl→3Galβl→4GlcNAcβl→3Galβl→4GlcNAcβl→O-(CH₂),CH₃ as acceptors (Gu et al., J. Biol. Chem. 267:2994-2999 (1992), which is incorporated herein by reference). Synthetic acceptors were kindly provided by Dr. Olé Hindsgaul, University of Alberta, Canada.

20 Results of these assays are shown in Table I.

Assuming transfection efficiency of the cells is approximately 20-30%, the level of enzymatic activity directed by cells transfected with pcDNAI-C2GnT is roughly equivalent to the level observed in HL-60 cells.

TABLE I

Core 2 $B1\rightarrow 6$ N-acetylglucosaminyltransferase activity in CHO-Py-leu cell extracts transfected with pcDNAI or pcDNAI-C2GnT.

5.				
	Vector	Core 2 ßl→6 GlcNAc transferase activity (pmol/mg of protein/hr)		
•	pcDNAI	n.d.		
10	pcDNAI-C2GnT	764		

CHO-Py-leu cells were transfected with pcDNAI or pcDNAI-C2GnT, as described in the specification. Endogenous activity was measured in the absence of acceptor and subtracted from values determined in the presence of added acceptor. Galβ1→3GalNAcα-p-nitrophenyl was used as an acceptor. n.d. = not detectable. For comparison, the core 2 β1→6 N-acetylglucosaminyltransferase activity measured in HL-60 cells under identical conditions was 3228 pmol/mg of protein per hr.

In order to unequivocally establish that C2GnT CDNA sequence encodes core B1→6 Nacetylglucosaminyltransferase, plasmid, pPROTA-C2GnT was constructed containing the DNA sequence encoding the putative catalytic domain of 2 core B1→6 Nacetylglucosaminyltransferase fused in frame with the signal peptide and IgG binding domain of S. aureus protein A (Fig. 7). The putative catalytic domain is contained in a 1330 bp fragment of the C2GnT cDNA that encodes amino 30 acid residues 38 to 428. Plasmid pPROTA was kindly provided by Dr. John B. Lowe.

The polymerase chain reaction (PCR) was used to insert EcoRI recognition sites on either side of the 1330 bp sequence in pcDNAI-C2GnT DNA. PCR was performed using the synthetic oligonucleotide primers 5'-TTTGAATTCCCCTGAATTTGTAAGTGTCAGACAC-3' (SEQ. ID. NO. 6) and 5'-TTTGAATTCGCAGAAACCATGCAGCTTCTCTGA-3' (SEQ. ID. NO. 7)

(EcoRI recognition sites underlined). The EcoRI sites allowed direct, in-frame insertion of the fragment into the unique EcoRI site of plasmid pPROTA (Sanchez-Lopez et al., <u>J. Biol. Chem.</u> 263:11892-11899 (1988), which is incorporated herein by reference).

The nucleotide sequence of the insert as well as the proper orientation were confirmed by DNA sequencing using the primers described above for cDNA sequencing. Plasmid pPROTA-C2GnT allows secretion of the fusion protein from transfected cells and binding of the secreted fusion protein by insolubilized immunoglobulins.

Either pPROTA or pPROTA-C2GnT was transfected into COS-1 cells. Following a 64 hr period to allow transient expression, cell supernatants were collected Cell (1990)). 15 (Kukowska-Latallo et al., supra, supernatants were cleared by centrifugation, adjusted to 0.05% Tween 20 and either assayed directly for core 2 β 1 \rightarrow 6 N-acetylglucosaminyltransferase activity or used in IgG-For the latter Sepharose (Pharmacia) binding studies. 20 assay, supernatants (10 ml) were incubated batchwise with approximately 300 μl of IgG-Sepharose for 4 hr at 4°C. The matrices were then extensively washed and used directly for glycosyltransferase assays.

No core 2 \$1 \to 6 N-acetylglucosaminyltransferase

25 activity was detected in the medium of COS-1 cells
transfected with the control plasmid, pPROTA. Similarly,
no enzymatic activity was associated with IgG-Sepharose
beads. In contrast, a significant level of core 2 \$1 \to 6 Nacetylglucosaminyltransferase activity was detected in the
30 medium of COS-1 cells transfected with pPROTA-C2GnT. The
activity also associated with the IgG-Sepharose beads
(Table II). No activity was detected in the supernatant
following incubation of the supernatant with IgG-Sepharose.

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TABLE II

Determination of Enzymatic Activities Directed by pPROTA-C2GnT.

5	Acceptors and linkages formed	Radioactivity (cpm) with (+) and without (-) acceptor		
		-	+	
10	GlcNAcB1 6 GalBl→3GalNAc (core 2-GnT)	109	1048	
15	GlcNAcB1 6 GlcNAcB1→3GalNAc (core 4-GnT)	111	113	
20	GlcNAcß1 6 GlcNAcß1→2Man (GnTV)	118	115	
25	GlcNAcB1 6 GlcNAcBl→3Gal (I-GnT)	111	113	
30	GlcNAcB1 6 Galßl→4GlcNAcßl→3Gal (I-GnT)	99	96	

COS-1 cells were transfected with pPROTA-C2GnT and the conditioned media were incubated with IgG-Sepharose. The proteins bound to the IgG-Sepharose were assayed for ß1→6 N-acetylglucosaminyltransferase activity by using appropriate acceptors. The linkages formed are indicated by italics. Similar results were obtained in three independent experiments.

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EXAMPLE VI

DETERMINATION OF C2GnT SPECIFICITY

Four types of ß1→6 Nacetylglucosaminyltransferase linkages have been reported,
including core 2 and core 4 in O-glycans, I-antigen and a
branch attached to mannose that forms tetraantennary Nglycans (see Table II). In order to determine whether
these different structures are also synthesized by the
cloned C2GnT cDNA sequence, enzymatic activity was
determined using five different acceptors.

As shown in Table II, the fusion protein was only active with the acceptor for core 2 formation. was true when the formation of $\mathfrak{S}1 \rightarrow 6$ N-acetylglucosaminyl linkage to internal galactose residues was examined (Table II, see structure at bottom). This result precludes the 15 likelihood that the enzyme encoded by the C2GnT cDNA sequence may add N-acetylglucosamine to a non-reducing terminal galactose. The HL-60 core 2 acetylglucosaminyltransferase is exclusively responsible for the formation of the GlcNAcβ1→6 branch on Galß1→3 20 GalNAc.

Although the invention has been described with reference to the disclosed embodiments, it should be understood that various modifications can be made without departing from the spirit of the invention. Accordingly, the invention is limited only by the following claims.

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SEQUENCE LISTING

- (1) GENERAL INFORMATION:
 - (i) APPLICANT: LA JOLLA CANCER RESEARCH FOUNDATION
 - (ii) TITLE OF INVENTION: A NOVEL BETA1-6 N-ACETYLGLUCOSAMINYLTRANSFERASE, ITS ACCEPTOR MOLECULE, LEUKOSIALIN AND A METHOD FOR CLONING PROTEINS HAVING ENZYMATIC ACTIVITY
 - (iii) NUMBER OF SEQUENCES: 8
 - (iv) CORRESPONDENCE ADDRESS:
 - (A) ADDRESSEE: CAMPBELL AND FLORES
 - (B) STREET: 4370 LA JOLLA VILLAGE DRIVE, SUITE 700
 - (C) CITY: SAN DIEGO
 - (D) STATE: CALIFORNIA
 - (E) COUNTRY: USA (F) ZIP: 92122

 - (V) COMPUTER READABLE FORM:
 - (A) MEDIUM TYPE: Floppy disk
 - (B) COMPUTER: IBM PC compatible
 - (C) OPERATING SYSTEM: PC-DOS/MS-DOS
 (D) SOFTWARE: PatentIn Release #1.0, Version #1.25
 - (vi) CURRENT APPLICATION DATA:
 - (A) APPLICATION NUMBER:
 - (B) FILING DATE: 30 September 1993
 - (C) CLASSIFICATION:
 - (viii) ATTORNEY/AGENT INFORMATION:
 - (A) NAME: KONSKI, ANTOINETTE F.
 - (B) REGISTRATION NUMBER: 34,202
 - (C) REFERENCE/DOCKET NUMBER: FP-LJ 9756
 - (ix) TELECOMMUNICATION INFORMATION:
 - (A) TELEPHONE: 619-535-9001
 - (B) TELEFAX: 619-535-8949
- (2) INFORMATION FOR SEQ ID NO:1:
 - (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 900 base pairs
 - (B) TYPE: nucleic acid
 - (C) STRANDEDNESS: both
 - (D) TOPOLOGY: linear
 - (ii) MOLECULE TYPE: DNA (genomic)
 - (ix) FEATURE:
 - (A) NAME/KEY: CDS
 - (B) LOCATION: 841..900
 - (ix) FEATURE:
 - (A) NAME/KEY: exon
 - (B) LOCATION: 91..192
 - (D) OTHER INFORMATION: /note= "EXON 1'IS LOCATED IN BOTH GENOMIC AND CDNA. IN THE CDNA EXON 1' IS IMMEDIATELY FOLLOWED BY EXON 2.

i	1 x	١ ١	FE	A'	TL	ΙR	E	:

- (ix) FEATURE:
 (A) NAME/KEY: exon
 (B) LOCATION: 359..428 (D) OTHER INFORMATION: /note= "EXON 1 IS LOCATED IN GENOMIC DNA"

(ix) FEATURE:

- (A) NAME/KEY: intron
 (B) LOCATION: 193..806
 (D) OTHER INFORMATION: /note= "THIS SEGMENT OF NUCLEIC ACID CONSTITUTES INTRON SEQUENCE OF THE CDNA"

(ix) FEATURE: .

- (A) NAME/KEY: exon
- (B) LOCATION: 807..900
- (D) OTHER INFORMATION: /note= "EXON 2 IS LOCATED IN BOTH GENOMIC AND CDNA. IN THE CDNA EXON 2 IMMEDIATELY FOLLOWS EXON 1'."

(xi) SEQUENCE DESCRIPTION: SEQ ID NO:1:

TTGGGGACCA CAAATGCAAA GGAAACCACC CTCCCCTCCC	60
GAGTTCTCAG GCTCACATTC CCACCACCCA CCTCTGAGCC CAGCCCTCCC TAGCATCACC	120
ACTTCCATCC CATTCCTCAG CCAAGAGCCA GGAATCCTGA TTCCAGATCC CACGCTTCCC	180
TGCCTCCCTC AGGTGAGCCC CAGACCCCCA GGCACCCCGC TGGCCCCTGA AGGAGCAGGT	240
GATGGTGCTG TCTTCGCCCA GCAGCTGTGG GAGCAGGCGG GTGGGGCAGG ATGGAGGGGT	300
GGGTGGGGTG GGTGGAGCCA GGGCCCACTT CCTTTCCCCT TGGGGCCCTG TCCTTCCCAG	360
TCTTGCCCCA GCCTCGGGAG GTGGTGGAGT GACCTGGCCC CAGTGCTGCG TCCTTATCAG	420
CCGAGCCGGT AAGAGGGTGA GACTTGGTGG GGTAGGGGCC TCAGTGGGCC TGGGAATGTG	480
CCTGTGGCTT GAAAAGACTC TGACAGGTTA TGATGGGAAG AGATTGGGAG CCATTGGGCT	540
GCACAGGGTC AGGGAGGCC AGGAGGGGCT GGTCACTGCT GGAATCTAAG CTGCTGAGGC	600
TGGAGGGAGC CTCAGGATGG GGCTGATGGG GGAGCTGCCA GCATCTGTTC CTCTGTCATT	660
TCTGATAACA GTAAAAGCCA GCATGGAAAA AACCGTTAAA CCGCAGGTTG GGCCTGGCCG	720
TTGGCAGGGA AGTGGGCAGA GGGGAGGCCC GGCCAGGTCC TCCGGCAACT CCCGCGTGTT	780
CTGCTTCTCC GGCTGCCCAC CTGCAGGTCC CAGCTCTTGC TCCTGCCTGT TTGCCTGGAA	840
ATG GCC ACG CTT CTC CTT CTC CTT GGG GTG CTG GTG G	888

GCT CTG GGG AGC Ala Leu Gly Ser

900

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									38							
(2)	INFO	RMAT	поп	POR	SEQ	ID 1	10:2:									
	(i) S	(A) (B)	LEN	GTH:	RACTE : 20 amino GY: 1	amir aci	o ac								
	(i	i) M	OLEC	ULE	TYPE	E: pr	rotei	.n								
	(x	i) S	EQUE	WCE	DESC	CRIPT	CION:	SEC) ID	No:	2:					
Met 1	Ala	Thr	Leu	Leu 5	Leu	Leu	Leu	Gly	Val 10	Leu	Val	Val	Ser	Pro 15	Aab	
Ala	Leu	Gly	ser 20	٠												
(2)	INFO	RMAT	NOI	FOR	SEQ	ID 1	10:3:									
	(i)	(A (B (C) LE) TY :) SI	NGTE PE: RAND	nucl	CTERI 105 h leic ESS: line	ase acid	pair l	s							
	(ii)	MOL	ECUL	E TY	PE:	CDNA	A									
	(ix)	(A) NA	ME/F		CDS 220.	150) 4								
	(ix)	(A) NA	ME/F		poly										
	(ix)	(A (B) NA	ME/F CATI HER	ON: INFO		314 TION:	/st		_	ame: MOQ					
	(xi)	SEQ	UENC	E DE	ESCR	IPTIC	ON: S	EQ 1	D NO	3:						
GTGA	AGTG	CT C	AGAA	TGGG	G C	AGGAT	rgtci	CCI	GGAI	ATCA	GCA	CTAAC	STG 1	ATTCA	AGACTT	60
TCCT	TACT	TT T	'AAA'	GTG	CT GO	CTCTT	CATI	TC	LAGA!	rgcc	GTT	GCAG	CTC :	IGAT <i>i</i>	AATGC	120
AAAC	TGAC	AA C	CTTC	AAG	ec ca	ACGA	CGGAC	GG.	AAA!	CAT	TGGT	rgcti	rgg 1	AGCAT	TAGAAG	180
ACTG	CCCT	IC A	СААА.	.GGA.I	AA TO	CCT	TTAE	A TTC	STTT					ACG I		234
CTG Leu	CGA A	AGG Arg	AGA Arg	CTT Leu 10	TTT Phe	TCT Ser	TAT Tyr	CCC Pro	ACC Thr 15	AAA Lys	TAC Tyr	TAC Tyr	TTT Phe	ATG Met 20	GTT Val	282
CTT Leu	GTT : Val 1	TTA Leu	TCC Ser 25	CTA Leu	ATC Ile	ACC Thr	TTC Phe	TCC Ser 30	GTT Val	TTA Leu	AGG Arg	ATT Ile	CAT His 35	CAA Gln	AAG Lys	330
CCT Pro	GAA : Glu 1	TTT Phe 40	GTA Val	AGT Ser	GTC Val	AGA Arg	CAC His	TTG Leu	GAG Glu	CTT Leu	GCT Ala	GGG Gly 50	GAG Glu	AAT Asn	CCT Pro	378

AGT Ser	AGT Ser 55	GAT Asp	ATT Ile	TAA neA	TGC Cys	ACC Thr 60	AAA Lys	GTT Val	TTA Leu	CAG Gln	GGT Gly 65	GAT Asp	GTA Val	TAA NeA	GAA Glu	426
ATC Ile 70	CAA Gln	AAG Lys	GTA Val	AAG Lys	CTT Leu 75	GAG Glu	ATC Ile	CTA Leu	ACA Thr	GTG Val 80	AAA Lys	TTT Phe	FÅ3 TYY	AAG Lys	CGC Arg 85	474
CCT Pro	CGG Arg	TGG Trp	ACA Thr	CCT Pro 90	Asp Asp	GAC Asp	TAT Tyr	ATA Ile	AAC Asn 95	ATG Met	ACC Thr	AGT Ser	GAC Asp	TGT Cys 100	TCT Ser	522
TCT Ser	TTC Phe	ATC Ile	AAG Lys 105	aga Arg	CGC Arg	TAU TYA	TAT Tyr	ATT Ile 110	GTA Val	GAA Glu	CCC Pro	CTT Leu	AGT Ser 115	AAA Lys	GAA Glu	570
GAG Glu	GCG Ala	GAG Glu 120	TTT Phe	CCA Pro	ATA Ile	GCA Ala	TAT Tyr 125	TCT Ser	ATA Ile	GTG Val	GTT Val	CAT His 130	CAC His	FA3 TA3	ATT Ile	618
GAA Glu	ATG Met 135	CTT Leu	GAC Asp	AGG Arg	CTG Leu	CTG Leu 140	AGG Arg	GCC Ala	ATC Ile	TAT Tyr	ATG Met 145	CCT Pro	CAG Gln	TAA Asn	TTC Phe	666
TAT Tyr 150	Сув	GTT Val	CAT His	GTG Val	GAC Asp 155	ACA Thr	AAA Lys	TCC Ser	GAG Glu	GAT Asp 160	TCC Ser	TAT Tyr	TTA Leu	GCT Ala	GCA Ala 165	714
GTG Val	ATG Met	GGC Gly	ATC Ile	GCT Ala 170	TCC Ser	TGT Cys	TTT Phe	AGT Ser	AAT Asn 175	GTC Val	TTT Phe	GTG Val	GCC Ala	AGC Ser 180	CGA Arg	762
TTG Leu	GAG Glu	AGT Ser	GTG Val 185	Val	TAT Tyr	GCA Ala	TCG Ser	TGG Trp 190	ser	CGG Arg	GTT Val	CAG Gln	GCT Ala 195	GAC Asp	CTC Leu	810
AAC Asn	тсс	ATG Met 200	Lys	GAT Asp	CTC Leu	тат туг	GCA Ala 205	Met	AGT Ser	GCA Ala	AAC Asn	TGG Trp 210	Lys	TAC Tyr	TTG Leu	858
ATA Ile	AAT Asn 215	Leu	тст Суз	GGT Gly	ATG Met	GAT Asp 220	Phe	CCC	ATT	Lys Lys	ACC Thi	AAC Asn	CTA Leu	GAA Glu	ATT Ile	906
GTC Val 230	Arg	AAG Lys	CTC Lev	AAG Lys	TTG Leu 235	TTA	ATG Met	GGA Gly	GAA Glu	AAC Asn 240	leA i	CTG Leu	GAA Glu	ACG Thr	GAG Glu 245	954
AGG	ATO	CCA Pro	TCC Ser	CAT His 250	Lys	GAA Glu	GAA Glu	AGG Arg	TGG Trp 255	Lys	AAC Lys	G CGG B Arg	TAT Tyr	GAG Glu 260	vai	1002
GT7 Val	IAA 1 18A 1	GG# Gly	AAC Lys	Leu	ACA Thr	DAA .	ACA Thi	GGG Gly 270	Thi	Val	C AAI L Ly:	A ATO	CT1 Leu 275	Pro	CCA Pro	1050
CT(Let	GA Glu	A ACI 1 Thi 280	r	CTC Lev	TTT Phe	TCT Ser	GG(Gl) 285	Sei	GCC Ala	TAC TY	r Ph	C GTC e Val 290	L Val	AGT Ser	AGG Arg	1098
GA(TA:	r Vai	G GGG	э тал ү туг	GTA Val	CTA Lev	Gli	AA E	GAI	A AA	TA A II e 0E	e Gl	A AAC	TTC	ATG Met	1146
GA Glu 31	u Tr	g GC	A CAI	A GAG	C ACA	Ту	AGC Set	c cc	r GA' o As _l	r GA p Gl 32	u Ty	T CTO	c TGG u Tr	G GCC P Ala	Thr 325	1194

ATC Ile	C AA Gln	AGG Arg	ATT Ile	CCT Pro 330	GAA Glu	GTC Val	ccg Pro	GGC Gly	TCA Ser 335	CTC Leu	CCT Pro	GCC Ala	AGC Ser	CAT His 340	AAG Lys	1242
TAT Tyr	GAT Asp	CTA Leu	TCT Ser 345	GAC Asp	ATG Met	CAA Gln	GCA Ala	GTT Val 350	GCC Ala	AGG Arg	TTT Phe	GTC Val	AAG Lys 355	TGG Trp	CAG Gln	1290
TAC Tyr	TTT Phe	GAG Glu 360	GGT Gly	GAT A sp	GTT Val	TCC Ser	AAG Lys 365	GGT Gly	GCT Ala	CCC Pro	TAC Tyr	CCG Pro 370	CCC Pro	TGC Cys	GAT Asp	1338
GGA Gly	GTC Val 375	CAT His	GTG Val	CGC Arg	TCA Ser	GTG Val 380	TGC Cys	ATT Ile	TTC Phe	GGA Gly	GCT Ala 385	GGT Gly	GAC A sp	TTG Leu	AAC Aan	1386
TGG Trp 390	ATG Met	CTG Leu	CGC Arg	TÀ2 YYY	CAC His 395	CAC His	TTG Leu	TTT Phe	GCC Ala	AAT Asn 400	AAG Lys	TTT Phe	gac A sp	GTG Val	GAT Asp 405	1434
GTT Val	GAC Asp	CTC Leu	TTT Phe	GCC Ala 410	ATC Ile	CAG Gln	TGT Cys	TTG Leu	GAT Asp 415	GAG Glu	CAT His	TTG Leu	AGA Arg	CAC His 420	AAA Lys	1482
GCT Ala	TTG Leu	GAG Glu	ACA Thr 425	TTA Leu	AAA Lys	CAC His	T GA	CCAT	TACG	GGC	TTAA	TTA	TGAA	ACAAC	IAA	1534
GAAG	GATA	CA C	AAA:	CGTA	C CI	TATO	TGTT	TCC	CCTI	CCT	TGTC	AGCG	TC C	GGAI	GATGG	1594
TATG	AAGT	CC I	CTTI	'GGGG	C AG	GGAC	TCTA	GTA	GATO	TTC	TTGT	'CAGA	GA A	AGCTO	CATGG	1654
TTTC	TGCA	GA G	CACA	GTTA	G CT	AGAA	AGGT	GAT	'AGCA	ATT.	AATG	TTCA	TC I	AGAC	TTAAT	1714
AGTG	GGAG	GA G	AAAT	.GGTA	G CC	TTGA	.GGCC	AGA	GCAG	GTA	GCAA	.GGCA	TT C	TGG <i>i</i>	AAGAG	1774
GGGA	CCAG	GG I	GGCI	'GGGG	A AG	AGGC	CGAT	GCA	AAAT.	.GTC	AGCC	TGTT	CC F	AGTO	CTCAG	1834
															LAATGT	1894
															AATCT	1954
															TAATT	2014
									GTCT	CCG	TATG	TCAT	CT C	AGGG	AGCTT	2074
'AAAA	TGGG	CT T	GATT	TAAC	A TT	GAAA	AAAA	A								2105

(2) INFORMATION FOR SEQ ID NO:4:

- (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 428 amino acids
 - (B) TYPE: amino acid
 (D) TOPOLOGY: linear
- (ii) MOLECULE TYPE: protein
- (xi) SEQUENCE DESCRIPTION: SEQ ID NO:4:

Met Leu Arg Thr Leu Leu Arg Arg Leu Phe Ser Tyr Pro Thr Lys

1 10 15

Tyr Tyr Phe Met Val Leu Val Leu Ser Leu Ile Thr Phe Ser Val Leu

Arg Ile His Gln Lys Pro Glu Phe Val Ser Val Arg His Leu Glu Leu Ala Gly Glu Asn Pro Ser Ser Asp Ile Asn Cys Thr Lys Val Leu Gln Gly Asp Val Asn Glu Ile Gln Lys Val Lys Leu Glu Ile Leu Thr Val Lys Phe Lys Lys Arg Pro Arg Trp Thr Pro Asp Asp Tyr Ile Asm Met Thr Ser Asp Cys Ser Ser Phe Ile Lys Arg Arg Lys Tyr Ile Val Glu Pro Leu Ser Lys Glu Glu Ala Glu Phe Pro Ile Ala Tyr Ser Ile Val Val His His Lys Ile Glu Met Leu Asp Arg Leu Leu Arg Ala Ile Tyr Met Pro Gln Asn Phe Tyr Cys Val His Val Asp Thr Lys Ser Glu Asp 145 150 160 Ser Tyr Leu Ala Ala Val Met Gly Ile Ala Ser Cys Phe Ser Asn Val Phe Val Ala Ser Arg Leu Glu Ser Val Val Tyr Ala Ser Trp Ser Arg 185 Val Gln Ala Asp Leu Asn Cys Met Lys Asp Leu Tyr Ala Met Ser Ala 200 Asn Trp Lys Tyr Leu Ile Asn Leu Cys Gly Met Asp Phe Pro Ile Lys Thr Asn Leu Glu Ile Val Arg Lys Leu Lys Leu Leu Met Gly Glu Asn Asn Leu Glu Thr Glu Arg Met Pro Ser His Lys Glu Glu Arg Trp Lys Lys Arg Tyr Glu Val Val Asn Gly Lys Leu Thr Asn Thr Gly Thr Val Lys Met Leu Pro Pro Leu Glu Thr Pro Leu Phe Ser Gly Ser Ala Tyr Phe Val Val Ser Arg Glu Tyr Val Gly Tyr Val Leu Gln Asn Glu Lys Ile Gln Lys Leu Met Glu Trp Ala Gln Asp Thr Tyr Ser Pro Asp Glu 315 Tyr Leu Trp Ala Thr Ile Gln Arg Ile Pro Glu Val Pro Gly Ser Leu Pro Ala Ser His Lys Tyr Asp Leu Ser Asp Met Gln Ala Val Ala Arg Phe Val Lys Trp Gln Tyr Phe Glu Gly Asp Val Ser Lys Gly Ala Pro Tyr Pro Pro Cys Asp Gly Val His Val Arg Ser Val Cys Ile Phe Gly 375

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42 Ala Gly Asp Leu Asn Trp Met Leu Arg Lys His His Leu Phe Ala Ask 385 390 Lys Phe Asp Val Asp Val Asp Leu Phe Ala Ile Gln Cys Leu Asp Glz His Leu Arg His Lys Ala Leu Glu Thr Leu Lys His 420 425 (2) INFORMATION FOR SEQ ID NO:5: (i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 34 base pairs (B) TYPE: nucleic acid (C) STRANDEDNESS: single (D) TOPOLOGY: linear (ii) MOLECULE TYPE: CDNA (xi) SEQUENCE DESCRIPTION: SEQ ID NO:5: TTTGAATTCC CCTGAATTTG TAAGTGTCAG ACAC 34 (2) INFORMATION FOR SEQ ID NO:6: (i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 33 base pairs (B) TYPE: nucleic acid (C) STRANDEDNESS: single (D) TOPOLOGY: linear (ii) MOLECULE TYPE: cDNA (xi) SEQUENCE DESCRIPTION: SEQ ID NO:6: TTTGAATTCG CAGAAACCAT GCAGCTTCTC TGA 33 (2) INFORMATION FOR SEQ ID NO:7: (i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 15 base pairs (B) TYPE: nucleic acid (C) STRANDEDNESS: single (D) TOPOLOGY: linear (ii) MOLECULE TYPE: protein (V) FRAGMENT TYPE: internal (ix) FEATURE: (A) NAME/KEY: CDS (B) LOCATION: 1..15 (D) OTHER INFORMATION: /note= "PROTEIN A - C2GNT FUSION PROTEIN"

(xi) SEQUENCE DESCRIPTION: SEQ ID NO:7:

GGG AAT TCC CCT GAA Gly Asn Ser Pro Glu 1 5

15

- (2) INFORMATION FOR SEQ ID NO:8:
 - (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 5 amino acids
 (B) TYPE: amino acid
 (D) TOPOLOGY: linear
 - (ii) MOLECULE TYPE: protein
 - (xi) SEQUENCE DESCRIPTION: SEQ ID NO:8:

Gly Asn Ser Pro Glu

CLAIMS

We claim:

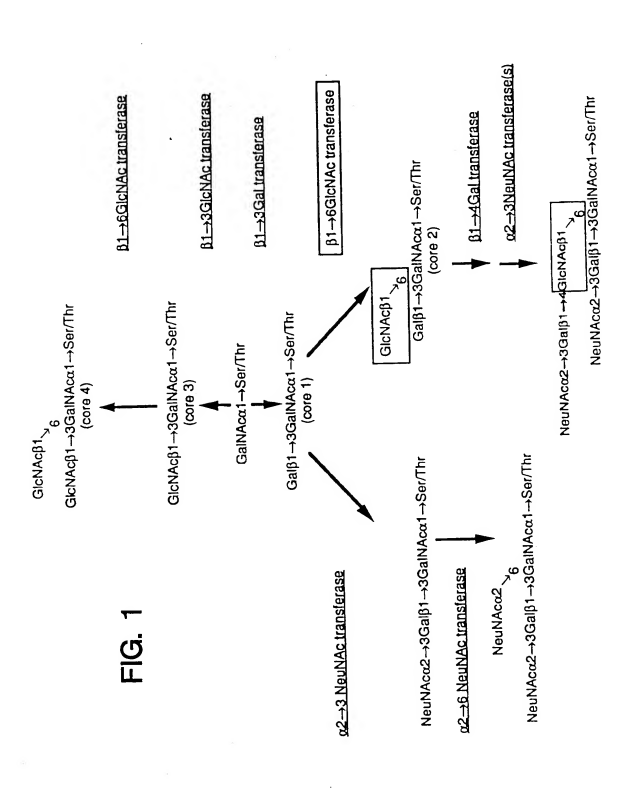
- 1. A purified human protein or an active fragment thereof having $\beta 1 \rightarrow 6$ N-5 acetylglucosaminyltransferase activity.
 - 2. The purified protein of claim 1, wherein said activity is that of UDP-GlcNAc:Galß1+3GalNAc (GlcNAc to GalNAc) ß1+6 N-acetylglucosaminyltransferase.
- 3. The purified protein of claim 2, wherein 10 said protein has a relative molecular weight of about 50 kD.
 - 4. An isolated nucleic acid encoding the human protein or active fragment thereof of claim 1.
- 5. A vector containing the nucleic acid of 15 claim 4.
 - 6. The vector of claim 5, wherein said vector is a plasmid.
 - 7. The vector of claim 5, wherein said vector is pcDNAI-C2GnT.
- 8. A host cell containing the vector of claim
 5.
 - 9. A purified human protein or a fragment thereof that is an acceptor molecule, said acceptor molecule being acted upon by the protein of claim 2 having activity which exclusively forms core 2 oligosaccharide structures in O-glycans.

- 10. The acceptor molecule of claim 9, wherein said acceptor molecule is leukosialin, CD43.
- 11. An isolated nucleic acid encoding the acceptor molecule of claim 9.
- 12. A vector containing the nucleic acid of claim 11.
- 13. The vector of claim 12, wherein said vector is a plasmid.
- 14. The vector of claim 12, wherein said vector is $pcDSR\alpha$ -leu.
- 15. A host cell containing the vector of claim 12.
- 16. A method of obtaining from a cell line, which does not normally contain a protein having catalytic activity or an acceptor molecule for said protein, a nucleic acid encoding said protein having catalytic activity comprising:
 - a. transfecting said cell line with a DNA sequence encoding the acceptor molecule, wherein the acceptor molecule is stably expressed in the cell line;
- b. transfecting said cell line with a cDNA 10 library containing said nucleic acid in a vector, wherein proteins encoded by the transfected cDNA are transiently expressed;
 - c. screening the transfected cells for expression of said protein having catalytic activity; and
- d. isolating the nucleic acid encoding the protein having catalytic activity.

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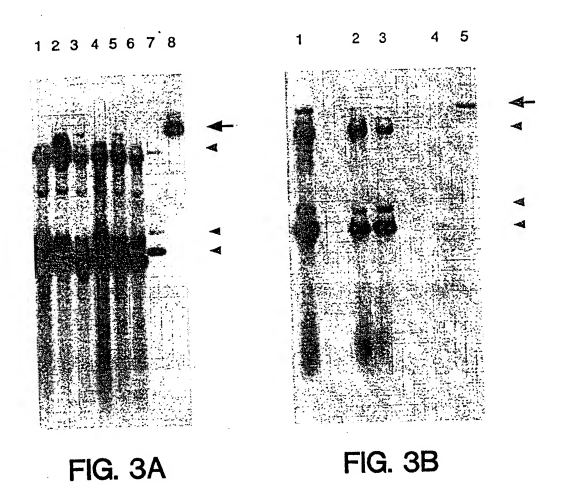
- 17. The vector of claim 16, wherein said vector replicates in the transfected cell line.
- 18. The vector in claim 17, wherein said vector is a plasmid.
- 19. The vector of claim 16, wherein said vector contains a viral replication origin.
- 20. The vector of claim 19, wherein said replication origin is the polyoma virus replication origin.
- 21. The cell line of claim 16, wherein said cell line supports replication of a vector.
- 22. The cell line of claim 16, wherein said cell line expresses polyoma virus large T antigen.
- 23. The cell line of claim 16, wherein said cell line is the Chinese hamster ovary cell line.
- 24. The cell line of claim 23, wherein said cell line is CHO-Py-leu.
- 25. A method of isolating a polypeptide having catalytic activity that forms core 2 oligosaccharide structures in O-glycans, said method comprising growing the host cell of claim 8 under conditions which favor expression of a nucleic acid encoding said polypeptide, and isolating said polypeptide so produced.

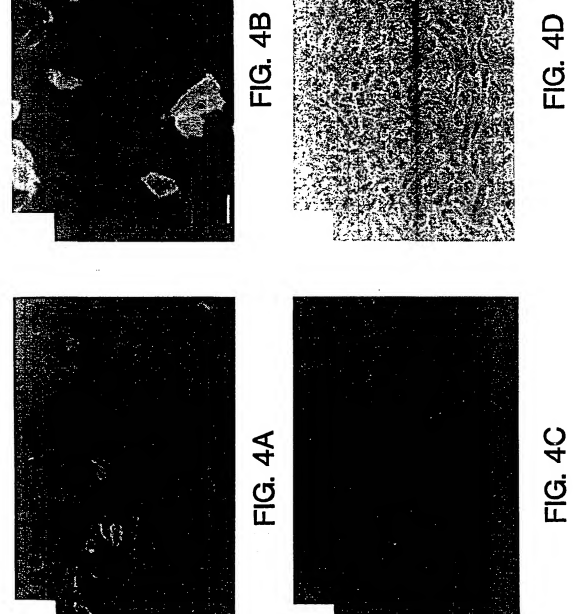


54 (20	ATGGCCACGCTTCTCCTTGGGGTGCTGGTGGTAAGCCCAGACGCTCTGGGGAGG	Exon2
48	CTGCTTCTCCGGCTGCCCACCTGCAGGTCCCAGCTCTTGCTCCTGCCTG	
42	TTGGCAGGGAAGTGGGCAGAGGGGCCCGGCCAGGTCCTCCGGCAACTCCCGCGTGTT	
36	TCTGATAACAGTAAAAGCCAGCATGGAAAAAACCGTTAAACCGCAGGTTGGGCCTGGCCG	
30	TGGAGGGAGCCTCAGGATGGGGCTGATGGGGGGGGGCTGCCAGCATCTGTTCCTCTGTCATT	
24	GCACAGGGTCAGGGAAGGCCAGGAGGGGCTGGTCACTGCTGGAATCTAAGCTGCTGAGGC	
18	CCTGTGGCTTGAAAAGACTCTGACAGGTTATGATGGGAAGAGATTGGGAGCCATTGGGCT	
12	CCGAGCCGGTAAGAGGGTGAGACTTGGTGGGGTAGGGGCCTCAGTGGGCCTGGGAATGTG	
9	TCTTGCCCCAGCCTCGGGAGGTGGTGGACCTGGCCCCAGTGCTGCGTCCTTATCAG	xon1
	GGGTGGGGTGGGGCCAGGGCCCACTTCCTTTCCCTTGGGGCCCTGTCCTTCCQAG	
- 5	GATGGTGCTGTCTTCGCCCAGCAGCTGTGGGAGCAGGCGGGGGGGG	
-11		
-179		xon1'
-23	GAGTICTCAGGCTCACATTCCCACCACCACCTCTGAGCCCAGCCCTCCCT	
-29	TTGGGGACCACAAATGCAAAGGAAACCACCCTCCCCTCC	

SUBSTITUTE SHEET

FIG. 2





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	-219 GTGAAGTGCTCAGAATGGGGCAGGATGTCACCTGGAATCAGCACTAAGTGATTCAGACTTTCCTTACTTTTAAATGTGCTGCTCTTCATTTCAAGATGC	-121
	CGTTGCAGCTCTGATAAATGCAAACTGACAACCTTCAAGGCCACGAGGGAAAATCATTGGTGCTTGGAGCATAGAAGACTGCCCTTCACAAAGGAAATCCCTGATTATTGTTTGAA	-
5	ATGCTGAGGACGTTGCTGCGAAGGAGACTTTTTCTTATCCCACCAAATACTACTTTATGGTTCTTGTTTTATCCCTAATCACCTTCTCCGTTTTAAGGATTCATCAAAAGCCTGAATTT M L R T L L R R R L F S Y P T K Y Y F M V L V L S L I T F S V L R I H Q K P E F	120
SUB	GTAAGIGICAGACACTIGGAGCIIGCIGGGGAGAAICCTAGIAGIGATAITAATIGCACCAAAGITITACAGGGIGATGTAAATGAAATCCAAAAGGTAAAGGTTGAGATCCTAACAGTG V S V R H L E L A G E N P S S D I N C T K V L Q G D V N E I Q K V K L E I L T V	240
STITE	AAATITAAAAAGGGCCCTCGGTGGACACCTGACGACTATATAAACATGACCAGTGACTGTTCTTTCT	360 120
JTE	TITCCAATAGCATATTCTATAGTGGTTCATCACAAGATTGAAATGCTTGACAGGCTGCTGAGGGCCATCTATAGCCTCAGAAATTGCGTTCATGGGACACAAAATCCGAGGAT	4 80 160
SHE	TCCTATITIAGCTGCAGTGATGGGCATCGCTTCCTGTTTTAGTAATGTCTTTGTGGCCGGATTGGAGGTGTGGGTTTATGCATCGTGGAGCCGGGTTCAGGCTGACTGCATG Isylaby Sylaby Siskon Color of Siskon Viron Siskon Siskon Siskon Siskon Siskon On Ciri	600
ET	A AGGATCTCTATGCAATGAGTGCAAACTGGAAGTACTTGATAAATCTTTGTGGTATGGATTTTCCCATTAAAACCAACC	720

FIG. 5A

	AACCIGGAAACGGAGGAIGCCAICCCATAAAGAAGAAGAAGAAGAAGAAGGGIAIGAGGICGIIAAIGGAAAACIGACAAAACAGAGGACIGICAAAAAIGCIICCICCACICGAAACA N L E T E R M P S H K E E R W K K R Y E V V N G K L T N T G T V K M L P P L E T	840 280
	CCTCTCTTTTCTGGCAGTGCCTACTTCGTGGTCAGTAGGGGTATGTGGGGTATGTACTACAGAATATCCAAAAGTTGATGGGGGGGG	960 320
SU	D TATCTCTGGGCCACCATCCAAAGGATTCCTGAAGTCCCGGGCTCACTCCCTGCCATAAGTATGATCTATCT	1080 360
BSTI	GOTGATGITICCAAGGGTGCTCCCTACCGCCCTGCGATGGAGTCCATGTGCGCTCAGTGTGCAGCTGGTGACTTGAACTGGATGCTGCGCAAACACCACTTGTTTGCCAAT	1200 400
TUT	AAGTITGACGTGGATGTTGACCTCTTTGCCATCCAGTGTTTGGATGACATTTGAGACACTTTGGAGACATTAAAAACACTGACCATTACGGGCAATTTTATGAACAAGAAGG	1320 428
E S	ATACACAAAACGTACCTTATCTGTTTCCCCTTCCTTGTCAGCGTCGGGAAGATGGTATGAAGTCCTCTTTGGGGCAGGGACTCTAGTAGATCTTCTTGTCAGAGAAGCTGCATGGTTTCT	1440
HE	GCAGAGCACAGTTAGCTAGAAAGGTGATAGCATTAAATGTTCATCTAGAGTTAATAGTGGGAGGAGTAAAGGTAGCCTTGAGGCCAGAGCAGGTAGCAAGGCATTGTGGAAAGAGGGGAC	1560
ET	CAGGGTGGCTGGGGAAGAGGCCGATGCATAAAGTCAGCCTGTTCCAAGTGCTCAGGGACTTAGCAAAATGAGAAGATGTGACCTGTGCCAAAACTATTTTGAGAATTTTAAATGTGACCA	1680
•	TITITCTGGTATG <u>AATAAA</u> CTTACAGCAACAAATAATCAAAGATACAATTAATCTGATATTTGTTGAAATAGAAATTTGATTGTACTATAAATGATTTTTGTAAATAATTTATAT	1800
	TCTGCTCTAATACTGTACTGTGTGTGTGTCTCCGTATGTCATCTCAGGGAGCTTAAAATGGGCTTGATTTAACATTGAAAAAAA	1886

FIG. SE

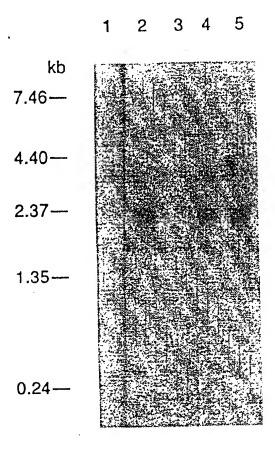
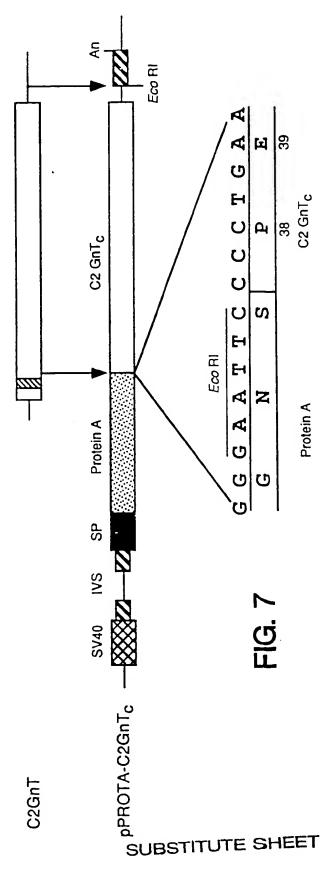


FIG. 6





INTERNATIONAL SEARCH REPORT

International application No. PCT/US93/09303

US CL.	Please See Extra Sheet. 435/6, 69.1, 193, 240.1, 252.3, 320.1; 530/395, 536	/23.2, 23.5							
	o International Patent Classification (IPC) or to both n	ational classification and IPC							
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1	ocumentation searched (classification system followed								
U.S. : 4	435/6, 69.1, 193, 240.1, 252.3, 320.1; 530/395, 536/	23.2, 23.3							
Documentat	ion searched other than minimum documentation to the	extent that such documents are included	in the fields scarched						
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Flasteraiad	ata base consulted during the international search (nar	ne of data base and, where practicable.	search terms used)						
	fline, Embase, Pascal, Derwent World Patent Index		,						
search ten	ms: c2gnt, acetylglucosaminyltransferase, core 2, gluc	osaminyltransferase							
	TO THE CONTRIBERED TO BE DELEVIANT								
C. DOC	UMENTS CONSIDERED TO BE RELEVANT								
Category*	Citation of document, with indication, where app	propriate, of the relevant passages	Relevant to claim No.						
x	Proceedings of the National Academy	of Sciences USA, Vol. 86,	9-15						
	issued February 1989, A. Pallant et al.,								
	encoding human leukosialin and localiz								
	to chromosomone 16", pages 1328-133	2, see the entire document.							
Y	Journal of Biological Chemistry, Vo	1. 266, No. 35, issued 15	1-8, 16-25						
1	December 1991, P. A. Ropp et								
	Purification and characterization								
	acetylglucosaminyltransferase", pages	23863-23871, see the entire							
	document.								
X Furth	ner documents are listed in the continuation of Box C	. See patent family annex.							
	ecial cutegories of cited documents:	"T" later document published after the int	emational filing date or priority						
.V. 90	cument defining the general state of the art which is not considered be part of particular relevance	date and not in conflict with the applic principle or theory underlying the in-	cation but cited to understand the						
	rlier document published on or after the international filing date	"X" document of particular relevance; the considered novel or cannot be considered.	ne claimed invention cannot be ered to involve an inventive step						
·L· do	cument which may throw doubts on priority claim(s) or which is of to establish the publication date of another citation or other	when the document is taken alone	La alaband insuration connect by						
ap-	ecial reason (as specified) cuspent referring to an oral disclosure, use, exhibition or other	"Y" document of particular relevance; if considered to involve an inventive combined with one or more other suc	e step when the document is						
204	2006	being obvious to a person skilled in the art							
tbe	cument published prior to the international filing date but later than priority date claimed	'&' document member of the same paten							
Date of the	actual completion of the international search	Date of mailing of the international se	aren report						
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PCT/US93/09303

Category*	Citation of document, with indication, where appropriate, of the relevant passages	Relevant to claim No.
Y	Blood, Vol. 77, No. 7, issued 01 April 1991, O. Saitoh et al., "T-lymphocytic leukemia expresses complex, branched O-linked oligosaccharides on a major sialoglycoprotein, leukosialin", pages 1491-1499, see the entire document.	1-8, 16-25
Y	Tetrahedon, Vol. 45, No. 17, issued 1989, E. J. Toone et al., "Enzyme-catalyzed synthesis of carbohydrates", pages 5365-5422, see pages 5409-5415.	1-8, 16-25
Y	Molecular and Cellular Biology, Vol. 11, No. 2, issued February 1991, D. F. Stern et al., "Spk1, a new kinase from Saccharomyces cerevisiae, phosphorylates proteins on serine, threonine, and tyrosine", pages 987-1001, see the entire document.	16-24
Y	Molecular and Cellular Biology, Vol. 8, No. 12, issued December 1988, S. Kornbluth et al., "Novel tyrosine kinase identified by phosphotyrosine antibody screening of eDNA libraries", pages 5541-5544, see the entire document.	16-24

INTERNATIONAL SEARCH REPORT

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C07K 13/00; C12N 1/21, 5/16, 9/10, 15/12, 15/54, 15/63, 15/79; C12P 21/00; C12Q 1/68									
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